A Cu\textsuperscript{I}/Cu\textsuperscript{III} prototypical organometallic mechanism for the deactivation of an active pincer-like Cu\textsuperscript{I} catalyst in Ullmann-type couplings†

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Unraveling the mechanistic details of copper-catalyzed arylation of nucleophiles (Ullmann-type couplings) is a very challenging task. It is a matter of intense debate whether it is a radical-based process or an organometallic redox-based process. The ancillary ligand choice in Ullmann-type couplings plays a key role in such transformations and can strongly influence the catalytic efficiency as well as the mechanism. Here, we show how a predesigned tridentate pincer-like catalyst undergoes a deactivation pathway through a Cu\textsuperscript{I}/Cu\textsuperscript{III} prototypical mechanism as demonstrated by helium-tagging infrared photodissociation (IRPD) spectroscopy and DFT studies, lending a strong support to the existence of an aryl–Cu\textsuperscript{III} species in the Ullmann couplings using this tridentate ligand.

Modern Cu-catalyzed cross-coupling reactions have recently evolved into reliable and efficient methods for the formation of C–C and C-heteroatom bonds, which are present in a large number of natural and pharmaceutical products.\textsuperscript{1–3} Unravelling the mechanistic details of copper-catalyzed arylation of nucleophiles (Ullmann-type couplings, Scheme 1a) is a very challenging task and a matter of intense debate in order to verify which of the two main mechanistic proposals is taking place: a radical-based or an organometallic-based process.\textsuperscript{4,5}

Since the late 1990s, much effort has been devoted to the use of chelating ligands, such as diamines, triamines, amino acids, phenanthroline derivatives and β-diketones, to perform coupling reactions under milder conditions while achieving enhanced yields.\textsuperscript{3,6,7} The detection of intermediate species after the activation of the aryl halide, which is usually rate-limiting, is very limited and most mechanistic proposals are derived from kinetic and computational studies.\textsuperscript{8–12} There is an ongoing discussion in the literature concerning the mechanistic pathway for Ullmann condensation reactions. The most invoked mechanisms for Ullmann couplings are based on either one-electron redox processes through radical intermediates that may operate via a Cu\textsuperscript{I}/Cu\textsuperscript{III} catalytic cycle, or two-electron redox processes via a Cu\textsuperscript{I}/Cu\textsuperscript{III} catalytic cycle (Scheme 1b).\textsuperscript{5} Recently, Peters, Fu and co-workers reported a photoluminescent Cu–carbazolide complex bearing monophosphine ligands for promoting the photoinduced C–N bond-forming reaction.\textsuperscript{13–15} Upon photoexcitation of the Cu–carbazolide complex, a Cu-containing radical is formed, as detected by EPR spectroscopy. This radical intermediate reacts with aryl halides via Single Electron Transfer (SET) to afford the corresponding C–N coupling product. Therefore, it is likely that several mechanisms might be possible simultaneously and the experimental conditions used are crucial to determine the operative one.

Focusing our attention on the design of appropriate auxiliary ligands for Ullmann couplings under standard thermal conditions,
we have previously demonstrated that a Cu(I)/Cu(III) catalytic cycle is operative in model macrocyclic aryl halide substrates, where active aryl-Cu(III) species have been completely characterized within these systems.\textsuperscript{16,17} Finding inspiration in the macrocyclic model systems, in this work we specifically design auxiliary ligand L\textsubscript{3} (Fig. 1) to reproduce the equivalent geometry of the Cu center, so that the stabilization of aryl-Cu(III) might become possible (Fig. 1).

In order to gain insight into the plausible operative mechanisms, we initially undertook a Cold-Spray\textsuperscript{18} MS study using L\textsubscript{3} as the auxiliary ligand for the C–O Ullmann-type coupling of iodobenzene and p-methoxyphenol as a model reaction (a standard 24 h reaction afforded a 45% yield of the diaryl ether product).\textsuperscript{19} Notably, an intense peak at \textit{m/z} = 304.0877 was detected in positive ESI mode after 0.5 hour (see the ESI for details of the procedure). At first sight, this peak could correspond to a putative Cu(III) intermediate species involved in the reaction ([L\textsubscript{3}–Cu(III)–(C\textsubscript{6}H\textsubscript{5})]\textsuperscript{+}), L\textsubscript{3} being the monodeprotonated version of L\textsubscript{3} (Fig. S1, ESI\textsuperscript{†}). We found the same analogous peak when using either bromobenzene (\textit{m/z} = 304.0877), 1-iodo-4-methylbenzene (\textit{m/z} = 318.1008) or 1-iodo-3,5-dimethylbenzene (332.1167) (Fig. S2–S4, ESI\textsuperscript{†}). Additionally, when using geometrically similar tridentate ligands L\textsubscript{4} (2,6-pyridinediyldimethanamine) and L\textsubscript{5} (N,N–diethyl-2,6-bis(aminomethylpyridine)), peaks analogous to 304.0877 were also observed at \textit{m/z} = 276.0540 and 332.1144, respectively (Fig. S5 and S6, ESI\textsuperscript{†}).

In an effort to validate whether this peak was a possible intermediate in the reaction mechanism, we extensively investigated the crude reaction mixture after 0.5 hour by characterizing the mass-selected ions at \textit{m/z} = 332.1167 formed when using L\textsubscript{3} and 1-iodo-3,5-dimethylbenzene (Fig. S4 in the ESI\textsuperscript{†}) by helium-tagging infrared photodissociation (IRPD) spectroscopy.\textsuperscript{20–22} IRPD spectroscopy provides well-resolved infrared spectra of mass-selected ions.\textsuperscript{23} The experimental IRPD spectrum of ions with \textit{m/z} 332 isolated from the crude reaction mixture after 0.5 h (a) compared to the theoretical IR spectra (B3LYP-D3/6-31G**+(Cu:6-311G*), scaling factor: 0.98) of other plausible species involved in the reaction (b–d).

When \textbf{[L\textsubscript{3}–Cu(III)]} was identified in the reaction mixture, we immediately considered the possibility that it could be the actual catalyst of C–O coupling. However, this was discarded based on our experimental findings. The use of the independently synthesized \textbf{[L\textsubscript{3}–Cu(III)]} complex as a catalyst (10 mol%) for the arylation of 2 afforded 16% yield of the biaryl ether product, compared to 45% yield when using \textbf{[L\textsubscript{3}–Cu(I)]} under the same catalytic conditions (Fig. S20, ESI\textsuperscript{†}). A similar drop in yield was found for the arylation of benzamide (from 75% down to 29%) and cyclohexylamine (from 21% down to 0%).

In light of these results, we propose that the Cu(I)/Cu(III) cycle may be the main operative mechanism of this coupling reaction, the detected species being the intramolecular reductive elimination product ([L\textsubscript{3}–Cu(III)] of the putative \textbf{[L\textsubscript{3}–Cu(III)–(C\textsubscript{6}H\textsubscript{5})]} species, \textit{i.e.} a decomposition product of the active catalyst. This non-radical mechanism may involve the formation of the \textbf{[L\textsubscript{3}–Cu(I)]} active species followed by a low-barrier aryl iodide oxidative addition.
and deprotonation of one of the secondary amines to produce \([L_3^{-}Cu^{II}-(aryl)]^+\). At this reaction crossroad, an axial ligand exchange by the nucelophile may trigger reductive elimination to afford the diaryl ether coupling product, or destabilization of the CuIII complex can induce intramolecular reductive elimination, yielding the Cu' amine-arylation by-product detected by MS.

DFT calculations were performed to unravel the mechanism of formation of the \([L_3^{-}CuH-(C_6H_5)]^+\) complex (Fig. 4a). The grouping of the aryl iodide adduct (B), which is coordinated in \(\eta^2\) fashion to the copper center \(d(Cu-C_{ipso}) = 1.999 \text{ Å}\) and \(d(Cu-C_{ortho}) = 2.045 \text{ Å}\), is endergonic \(\Delta G = 5.8 \text{ kcal mol}^{-1}\). However, the B adduct easily undergoes oxidative addition to the \([L_3^{-}Cu^{III}-(C_6H_5)]^+\) species (D) through the transition state C, which is only 1.6 kcal mol\(^{-1}\) higher in energy than B \((i.e. \Delta G^{\ddagger} = 7.4 \text{ kcal mol}^{-1})\). Removal of an iodine anion and NH deprotonation of D affords the \([L_3^{-}Cu^{III}-(C_6H_5)]^+\) intermediate (E). Finally, E proceeds to the detected species \([L_3^{-}Cu-(C_6H_5)]^+\) (G) through the reductive elimination transition state F with a barrier of 10.5 kcal mol\(^{-1}\). Transition state F involves a conformational change of the aryl ligand to achieve a proper geometrical orientation that favours the aryl-N coupling. Precedents of arylation of amines are known using well-defined macrocyclic aryl-CuIII model systems and standard Ullmann systems. DFT calculations predict that the geometry adopted by the copper centre significantly varies depending on the oxidation state of the metal (Fig. S18 and S19, ESI†). Further experimental substantiation of this structural variability is provided by the crystal structures of the tetrahedral \([L_3Cu^0(OTf)]_n\) and square-pyramidal (tetragonally distorted) \([L_3Cu^{III}(OTf)]_2[OTf]\) complexes shown in Fig. S18 and S19 (ESI†).

In the presence of \(p\)-methoxyphenolate, theoretical calculations predict a favourable aryl-O coupling to form the biaxial ether product (with a free energy barrier of 8.6 kcal mol\(^{-1}\)). The competing aryl-N coupling leading to catalyst decomposition proceeds via a higher energy barrier (11.7 kcal mol\(^{-1}\); Fig. 4b).

The presence of phenolate makes the aryl-N coupling less favourable, because it turns nitrogen deprotonation, \(i.e.\) the step preceding C-N coupling, from an exergonic to an endergonic process \((-2.3 \text{ kcal mol}^{-1}\) for the D \(\rightarrow\) E transformation in the absence of phenolate vs. 5.0 kcal mol\(^{-1}\) for the H \(\rightarrow\) K transformation when phenolate is present; compare Fig. 4a and b). After amine deprotonation, K first undergoes the required conformational change of the aryl ligand that has a free energy cost of 6.7 kcal mol\(^{-1}\) (M). Finally, the very reactive intermediate M proceeds to the detected species O through a barrierless reductive elimination transition state (N). Mechanistic insights on the selectivity between C-O and C-N couplings in Ullmann catalysis have been documented.

Therefore, a Cu'CuIII mechanism underlying the coupling reaction of iodobenzene and phenol derivatives stems from the above experimental and theoretical data; further proof was obtained by conducting the coupling experiment using the radical clock 1-allyloxy-2-iodobenzene (re) as the substrate and \(p\)-methoxophenol (2), benzamide (3) and cyclohexamine (4) as nucleophiles (see Scheme 2). The formation of the...
cylized coupling compounds (re-cyc-x) was not detected in any reaction, thus indicating the unfeasibility of a radical mechanism. We did observe relevant amounts of compound re-H (up to 25% with cyclohexylamine 4 as a nucleophile), where the iodine atom has been substituted by an H atom. Protodecupuration of a putative organometallic aryl–Cu bond has been already observed in a recent report.19

In conclusion, the efficient auxiliary ligand \( L_3 \) in C–O \( \text{Ullmann-type couplings} \) undergoes a decomposition pathway following a Cu\(^{II}\)/Cu\(^{III}\) mechanism via intramolecular arylation of one of the secondary amines of the complex. The helium tagging IRPD spectroscopy and DFT mechanistic studies, along with the absence of cyclized products using an \( r_c \) radical clock and the observation of protodecupuration products, strongly support the existence of an aryl–Cu\(^{III}\) species in Ullmann couplings using this specially designed tridentate \( L_3 \) ligand. Insights on decomposition gained from this model system has turned out to be valuable in catalyst and process design work, shedding light on the complex chemistry of Ullmann couplings.

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Notes and references